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Assessing Lignin Decomposition and Soil Organic Carbon Contents Across a Tropical Savannah-Rainforest Boundary in Guyana

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The soils beneath the rainforest of Guyana have the potential to hold, and release, large stores of carbon under land use and climate change. Little is known about soil carbon stocks or molecular dynamics in this region. This study therefore aims to elucidate differences in the molecular (lignin and tannin) and bulk soil organic carbon (SOC) stocks in different 'sub-environments' along a rainforest-savannah boundary, setting a framework for further investigation into the soil carbon dynamics of the region. Bulk SOC analysis shows that Gleysols have the highest stocks, particularly those under rainforest vegetation (swamp and island forests surrounded by savannah), whereas Plinthosols have significantly lower SOC stocks. Texture and soil water content analysis indicates that predominantly clay soils play a role in high SOC stocks, whilst predominantly sandy soils prevent SOC stocks from accumulating. Clay and sand are present in both Gleysols and Plinthosols, to different extents. Analysis of lignin and tannin in surface soils of the sub-environments reveals clear differences in molecular composition. Heavily degraded lignin signatures in rainforest Gleysols suggests a surrounding physio-chemical environment which promotes their degradation. Conversely, Plinthosols beneath woodland within the savannah have the greatest amount of lignin and tannin products. The presence of the clay mineral kaolinite and iron oxide strengite in these soils indicates a low ability for protection or complexing of organic matter. Therefore, water content and microbial activity may play a more important role in the degradation of lignin and tannin, as well as the SOC stock. With the potential for future deforestation due to land use or climate change, the high lignin degradation of Gleysols indicates a vulnerability to savannah encroachment. Forest Islands isolated from the main forest biome are the most vulnerable to change, and could lose a significant proportion of their SOC stock in a transition to savannah.

Keywords: lignin, tannin, thermochemolysis, ¹³C-labeled TMAH, soil organic carbon, savannah-rainforest boundary, Amazon, climate change

INTRODUCTION

Soil organic matter (SOM) is a key component of carbon storage in natural ecosystems. However, pressures from land use change, for example from forest to agriculture, and climate change altering the local weather patterns potentially impacts the amount of carbon stored in SOM through exposing it to greater degradation, increasing CO₂ emissions from the soil (Lal, 2004). Amazonian soils store approximately 66.9 Pg C in the top meter, accounting for around 4% of total global carbon stocks (Batjes and Dijkshoorn, 1999). Whilst some areas of the Amazon, which have large potentials to store carbon, have experienced extensive deforestation for agriculture, Guyana's rainforest remains largely intact. They cover 85% of the land and are bordered by the Rupununi savannah close to Brazil (Palo, 1994). It is therefore likely that soils under rainforest in Guyana hold large carbon stores, however there is less certainty of these stores on the savannah boundary.

In this study we focus on lignin, which is a key component of SOM, constituting about a third to the organic carbon content of the biosphere (Boerjan et al., 2003). Our knowledge of lignin stability and degradation in tropical settings is limited mainly to agricultural systems (Guggenberger et al., 1995; Olk et al., 2002; Filley et al., 2006; Rumpel et al., 2007), whereas little is known about lignin stability and degradation in natural tropical systems. There are some examples of lignin degradation in tropical ecosystems, although these are mainly in wetlands (Bala Krishna Prasad and Ramanathan, 2009; Spencer et al., 2010; DeAngelis et al., 2011).

Soil organic matter stores carbon in complex compounds, for instance lignin, and releases carbon as CO₂ primarily through microbial decomposition but also through abiotic factors. In woody plants, such as those found in rainforest and savannah systems (less frequent in the latter), lignin can contribute 30% to mass, whilst non-woody plants, including agricultural crops, show lesser amounts (Adler, 1977). Lignin has previously been considered recalcitrant and therefore preserved in soil due to the complexity and variability in linkages within the lignin biopolymer and its high molecular weight (R. L. Crawford, 1981). However, this recalcitrance has been contested and it is now known that its stability and preservation within soils is largely due to the physio-chemical environment surrounding it (Thevenot et al., 2010). Lignin degradation is fastest under aerobic conditions (Vane and Abbott, 1999; Kiem and Kögel-Knabner, 2003; Dignac et al., 2005; Bahri et al., 2006; Kalbitz et al., 2006; Marschner et al., 2008; Klotzbücher et al., 2011; Dungait et al., 2012), where white rot fungi are one of the most efficient degraders (Gold et al., 1989; Hatakka, 1994; Robertson et al., 2008). Some species of bacteria (*Streptomyces* and *Nocardia*) have also been found to degrade lignin (Sørensen, 1962; Antai and Crawford, 1981; Crawford et al., 1983; Godden et al., 1992; Trojanowski, 2001). In addition to biotic degradation, abiotic factors also play a role in the breakdown of lignin, for example through photo-oxidation (Hernes and Benner, 2003; Gallo et al., 2006; Frouz et al., 2011) and leaching (Hernes et al., 2007).

The rainforest and savannah regions in Guyana support a range of different vegetation types. These are likely to affect SOM

stocks and cycling through variations in leaf and wood litter chemistry, especially the abundance and type of lignin, tannin and carbohydrates present (Carr et al., 2013). Similarly, climate change may affect the ability of these soils to act as a carbon store. Currently, the seasonality of these ecosystems is divided between two main weather patterns; a coastal 'Tropical Wet' (or 'Tropical Rainforest') climatic region of two wet and two dry seasons and a 'Tropical Wet-Dry' (or 'Savannah') climatic region further inland covering the Guyana savannah, where a single wet and dry season prevails, therefore providing a smaller total precipitation than the 'Tropical Wet' pattern (Bovolo et al., 2012). However, with climate and land use change there is a potential for these patterns to change toward overall drier and hotter conditions (McSweeney et al., 2010). Hotter conditions are expected to increase the rate at which bacteria can break down organic carbon by heterotrophic respiration, speeding up the rate of chemical reactions (Lloyd and Taylor, 1994; Fenner and Freeman, 2011; Wang et al., 2014). Future alterations in hydrology are less certain than those in temperatures, but hydrology often determines the redox potential of soils, impacting oxic decomposition, with a decrease in microbial decomposition when flooded and intensive degradation during dryer seasons, as microbes utilize moisture remaining from the wet season (Fenner and Freeman, 2011; Bovolo et al., 2012).

Studies that have assessed molecular SOM in natural systems show that vegetation types have differing molecular carbon inputs into soils, which may be either more or less stable and can alter the SOC stock when these vegetation types shift (Guggenberger et al., 1995; Filley et al., 2006; Stewart et al., 2011; Carr et al., 2013). More recent studies have detected that polysaccharides, previously thought of as labile, are a major pyrolysis-GC/MS product in cultivated dystric cambisols, thus actually relatively stable (Gleixner et al., 2002). It has also been suggested that lignin has a faster turnover rate than bulk SOM in some temperate zone grassland and arable soils (Dignac et al., 2005; Heim and Schmidt, 2007). This is contrasted by other studies observing that lignin is not always preferentially degraded (Van Bergen et al., 1997; Kiem and Kögel-Knabner, 2003; Nierop et al., 2003).

In order to better understand the fate of lignin in pristine tropical ecosystems, we have performed a detailed biogeochemical study in a largely natural ecosystem in the Northern Rupununi, Central Guyana. A reconnaissance survey of soils and vegetation over a c. 10 km section of savannah-rainforest boundary was preliminarily undertaken. Following this, a representative section for a detailed transect study of bulk and molecular carbon analysis was chosen, running 2.5 km either side of this boundary. This paper presents an investigation of the molecular characteristics of the SOM and the SOC stock along this boundary. The surface soil is the most critical zone in the profile for carbon dynamics, particularly on the molecular scale, as it is where the majority of dead plant material (above and below ground) is deposited and where most microbial activity occurs (Foster, 1988). Further, lignin and tannin phenols have been shown to be most abundant in the surface soil, decreasing significantly with depth through the profile (Nierop and Filley, 2007; Mason et al., 2009). We therefore semi-quantitatively assess the amount of lignin and tannin products from samples

of living plant tissues, freshly deposited litter and surface soil (0–6 cm) and their extents of degradation using thermally assisted hydrolysis and methylation (THM) in the presence of tetramethylammonium hydroxide (TMAH) (Mason et al., 2009).

The aim of the study is to elucidate differences in the composition of molecular SOM and size of the SOC stock under a mosaic of vegetation types on the savannah-rainforest boundary. This baseline data collection and the differences observed in molecular and bulk SOC in the study region provide a framework for further investigation relating to the soil dynamics and their potential alteration through changes in seasonality and impact from climate extreme events (droughts/floods), both linked to climate change. This study provides critical geochemical data across sub-settings of a savannah-forest boundary that is commonly observed in the Amazon system. We present data from soil samples that have been obtained during the end of the dry season 2012, leaving studies that resolve seasonal or multi-annual variability still to be done.

We hypothesize that vegetation will have a dominant effect on the SOM composition and that the soil type will have a strong influence on the degradation state of this SOM. Consistent with other studies (Swain et al., 2010; Angst et al., 2021) we postulate that SOM degradation is inversely related to SOC stock. SOM degradation, defined as the break-down of organic matter in the soil, is controlled by the interplay of climate (precipitation and temperature), vegetation type and soil matrix factors including the physio-chemical environment and microbes. In order to achieve this aim, the objectives of this study were to:

- (a) Quantify the surface (0–6 cm) and full profile SOC stock of undisturbed soils across a savannah-rainforest transect.
- (b) Identify the molecular chemical composition (lignin and tannin) of vegetation inputs and surface soils across the transect.
- (c) Investigate the relationship between the extent of lignin degradation and the SOC stock.
- (d) Investigate how variations in soil texture, moisture and vegetation may impact SOC stocks and lignin degradation.

MATERIALS AND METHODS

Study Site

The area of this study is within the Northern Rupununi region of central Guyana. The Rupununi region is situated in the north eastern part of the Takutu Basin – a Mesozoic graben (280 km long and 40 km wide) between parallel faults which form the Pakaraima Mountains to the north and Kanuku Mountains to the south. The study sites are within the tribal lands of Wowetta at 4°01'02.5"N, 59°04'06.0"W (Figure 1). Following a reconnaissance survey of an area covering 10 km along the rainforest-savannah boundary, a transect of the most representative soils and vegetation was identified for sampling. From the boundary, the transect extends 2.5 km into the rainforest and 2.5 km into the savannah. Along this transect the topography undulated through flat areas, small hills or mounds of hardened laterite and wet depressions. The sites along

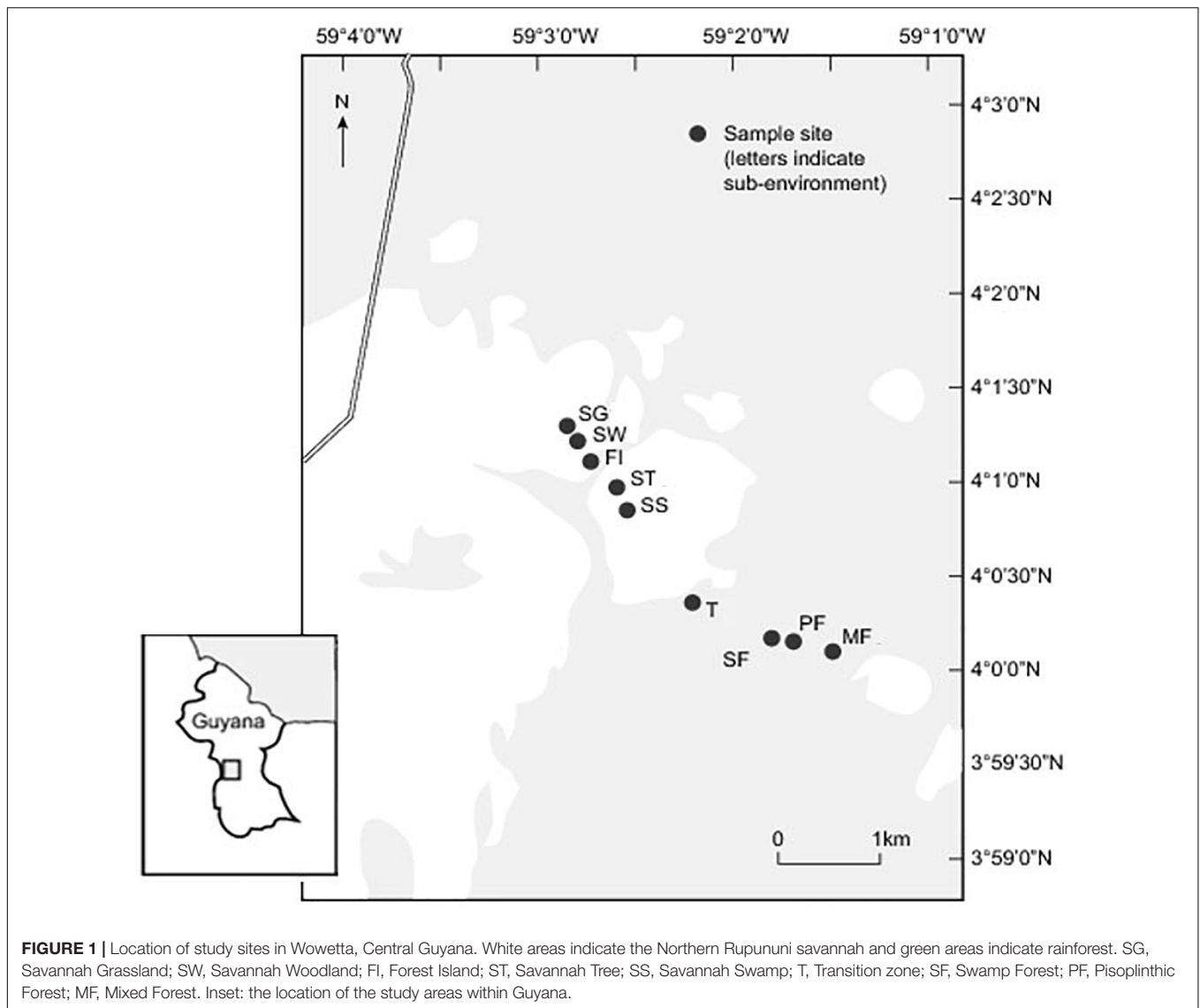
this transect were named Savannah Grassland (short grasses with no trees), Savannah Woodland (clumps of trees forming small woods within the predominantly grass-covered savannah), Savannah Tree (an isolated tree within the grassy savannah), Savannah Swamp (a waterlogged area of grasses, sedges and scattered palm trees which are few in number), Forest Island (an area of rainforest trees surrounded by savannah), Transition zone (dense scrub vegetation directly between the savannah and rainforest proper), Swamp Forest (a waterlogged topographical depression within the rainforest), Pisoplinthic Forest (an area of mixed rainforest tree spp. on a mound of hardened lateritic nodules – pisoplinthic material) and Mixed Forest (an area of level ground with mixed rainforest tree spp.) We use the term 'sub-environment' to refer to these sites from this point onward. Similar vegetation types and undulating landscape topography have been found in previous studies of the area (Eden, 1964; Sinha, 1968; Daniel and Hons, 1984; ter Steege, 2000). Although the area of savannah in Guyana is relatively small, its pristine quality provides a unique opportunity to investigate this at a climate sensitive interface (Bovolo et al., 2012).

Sample Collection and Preparation

Nine sample pits were excavated in February 2012, at the end of the main annual dry season, in order to sample and assess soils in advance of upcoming wet season floods. During sample collection, soil profile descriptions were taken and recorded according to the FAO (IUSS Working Group WRB, 2015). These included soil texture ('hand texture method'), soil color (Munsell Color chart) structure, boundary depth of each horizon and full sampling depth of the soil profile. Water table depth from the surface was also recorded using measuring tape as soon as the water table had been breached.

Biomass samples (fresh shoot/leaf litter and wood litter) were sampled in triplicate from the surface of the soil pit and the surrounding area (within 0.25 m of the soil pit). Where no litter layer existed, such as in the savannah grassland sites, fresh shoots were sampled. Fresh grass shoots were also sampled in areas of savannah with trees in addition to leaf litter, as the grass itself did not form a litter layer. All soil samples (including for bulk density) were taken in triplicate from 0–6 cm from the surface of each soil profile as well as at the mid-point between the top and bottom of each horizon in every soil profile. Bulk density sampling excluded the pisoplinthic forest (PF), where the pisoplinthic material (hardened aggregates of iron oxide) prevented accurate sampling. Volumetric water content (θ) was calculated using the weight of the pre-dried soil, the weight of oven dry soil, the known volume of bulk density sample and the bulk density (Rowell, 1994). It is important to note that water content was only measured at one point in time, from each triplicate of soil.

All samples for molecular analysis were stored in sealed amber glass jars and stored at -20°C from the field to the home laboratory to avoid any out of *situ* degradative changes. An aliquot of all samples was taken in order to conduct pH and molecular organic matter analysis on. Samples were then freeze dried and ground to a fine fraction ($<0.25\ \mu\text{m}$) and passed through a $<0.25\ \mu\text{m}$ sieve to verify the particle size.



Subsequently, organic soil compounds were extracted using Accelerated Solvent Extraction (ASE).

X-ray diffraction was undertaken on a subset of surface (0–6 cm) soil samples across the transect, covering the range of soil types found (including one sample from the Swamp Forest, Mixed Forest, Transition zone, Savannah Swamp, Savannah Woodland and Savannah Tree soils). Surface samples were selected to correspond to the molecular analysis also conducted at the surface of soil profiles (see Section “THM with TMAH analysis” below), in order to better understand the abiotic factors which may affect the state of molecular degradation. For example, the mineralogy may have consequences for the ability of the soil to protect or complex with organic matter (Gu et al., 1994; Sollins et al., 1996; Arnarson and Keil, 2000; Lützow et al., 2006). X-ray diffraction was carried out using a PANalytical X’Pert Pro Multipurpose Diffractometer (MPD) fitted with an X’Celerator detector, using 500 mg aliquot of the dried, ground and sieved (<0.25 μm) soil samples.

Soil Organic Carbon Concentrations and Stocks

Total Organic Carbon (TOC) concentration was determined using 0.1 g of freeze dried, sieved (to < 0.25 μm) and ground sample. Hydrochloric acid of 4.0 mol/L was added at an amount of 1 mL to each crucible and was drained from the sample for 4 h. The samples were then placed in the oven at 60 to 70°C for 16–24 h. A blank was used with each sample set.

After calculating both TOC concentration and bulk density it was then possible to calculate the weight of organic carbon (t C ha⁻¹) of soils by horizon, using the following formula:

Equation 1 Soil organic carbon stock (SOC):

$$SOC = d \times Db \times TOC$$

Where d = depth (cm), Db = bulk density (g cm³), TOC = total organic carbon (%).

Through using TOC concentration as a percentage, the conversion from g per cm³ to t per ha, is avoided, which has otherwise caused confusion and errors in the past (Cannell and Milne, 1995).

Where horizons were missing bulk density data, due to an inability to sample them (e.g., water saturation or depth of horizon), bulk densities were estimated using surrogate values determined by pedotransfer functions or rules (Bouma and Van Lanen, 1987), as used in Batjes and Dijkshoorn (1999). The primary surrogate option was to calculate the mean bulk density of the soil profile and use this value for missing horizons within the profile. The secondary option (if only one bulk density value was available for a profile) was to use a mean soil profile bulk density value of a corresponding soil type and the one available horizon. These surrogate values have been indicated using a “*” in **Table 1** below. The Pisoplinthic Forest soil did not have any similar profiles to use as surrogates, due to the hard lateritic nodules that it consisted of, therefore SOC stock for this profile has not been estimated.

THM With TMAH Analysis

Lignin and tannin phenols of vegetation and surface soil were assessed using on-line thermally assisted hydrolysis and methylation (THM) in the presence of unlabeled and ¹³C-labeled tetramethylammonium hydroxide (TMAH) (Mason et al., 2009). For the standard, 5 α -androstane with dichloromethane was used. THM was performed using a pulsed mode open pyrolysis system specifically a CDS 1000 pyroprobe unit (Chemical Data Systems, United States) fitted with a platinum coil and a CDS 1500 valved interface, as previously published (Mason et al., 2009).

The use of ¹³C-labeled TMAH deconvolutes the contribution of intact lignin phenols [guaiacyl (G), cinnamyl (C), and syringyl (S)] from microbially altered lignin or non-lignin sources including tannins (Filley et al., 1999). For example, the unlabeled thermochemolysis product S6 (3,4,5-trimethoxy benzoic acid methyl ester) is often a composite of syringic acid (lignin-derived) and gallic acid (tannin-derived) with the same mass spectra, however this will have different mass spectra following ¹³C-labeling (Mason et al., 2009). Thus, the amounts of unlabeled products can be corrected to give the “true” quantities of intact lignin present enabling accurate calculation of the lignin phenol parameters, e.g., lambda (Λ) (Filley et al., 2006; Nierop and Filley, 2007; Mason et al., 2009). Individual lignin phenols were corrected via the calculation of the hydroxyl contents of intact lignin, altered demethylated lignin and non-lignin sources. Source indicators C/G and S/G are used for non-woody and angiosperm/gymnosperm inputs, calculated using lignin phenols as (P18 + G18)/(G4 + G6) and (S4 + S6)/(G4 + G6), respectively (Filley et al., 2006). Ratios of the acid (Ad) and aldehyde (Al) endmembers of the lignin phenols guaiacyl and syringyl, as represented by [Ad/Al]_G and [Ad/Al]_S (calculated as G6/G4 and S6/S4) indicate the level of oxidation for guaiacyl and syringyl phenols, respectively (Filley et al., 2006). Increasing [Ad/Al]_{G/S} values indicate increasing oxidation, therefore, it is usually expected that these ratios are higher in soils than vegetation (Filley et al., 1999). Lambda (Λ) was calculated

as the sum of amounts of the 8 dominant lignin phenols (G4 + G5 + G6 + S4 + S5 + S6 + G18 + P18) (Filley et al., 2006).

In order to determine gallic acid (tannin) and demethylated lignin sources, the % aromatic hydroxyl contents of S6 (1% OH, 2% OH, and 3% OH) were determined using ¹³C-labeled TMAH. 1% OH refers to intact lignin, 2% OH to microbially demethylated lignin (syringic acid) and 3% OH to gallic acid (tannin derived). Filley et al. (2006) showed how multiple sources such as intact lignin and microbially altered lignin (specifically their methoxyl functionalities) can lead to the formation of identical products. Using previously established equations (Filley et al., 2006), these hydroxyl contents were calculated and then converted into mg/100 mg OC. The unlabeled TMAH thermochemolysis of these samples allowed the identification of S6, and thus the baseline for accurately calculating the ¹³C % addition.

RESULTS

Soil Classifications

The soil profiles sampled consist of Plinthosols and Gleysols with a range of sub-classifications and an underlying presence of iron oxide in plinthic form throughout the transect (**Figure 2**; FAO-ISRIC, 2006). In albic Plinthosols at SG, SW and ST sites, TOC concentration ranges from 0.03 to 1% and pH from 4.3–5.7 with greater acidity and lower SOC in the surface 6 cm. The color of this horizon changes from dark grayish brown in the SG to dark brown/very dark grayish brown beneath SW and ST, likely caused by different vegetation inputs or a persistence of SOM in the latter sites (see **Figures 6C,D** depicting more degraded lignin phenols which may indicate older SOM and **Figure 7** which shows significantly greater gallic acid in SW and ST than SG). Hand texture assessments confirm predominantly sandy textures (sand – sandy clay loam, generally increasing in clay with depth). The SW has a sandy loam surface soil compared to the sand texture of ST and SG, therefore, a slightly high clay content, however the water content, which may therefore also be greater and result in less degradation of organic carbon, does not differ between SW and ST soils. MF is an acric Plinthosol; the surface soil is pH 4.3, has a TOC concentration of 1.1% and a silt loam surface texture grading to clay with depth. PF is a pisoplinthic Plinthosol with a pH of 4.5, a TOC concentration of 2.7% and a mix of pisoplinthic material and coarse sandy texture. Sites FI and SF are both humic Gleysols, with black humified surface horizons that have high TOC concentrations, the FI average being roughly half of SF (6% and 13% respectively). The texture here is predominantly clay, although some sand does exist (e.g., the main profile texture of SF is sandy clay). SS and T soils are plinthic Gleysols with pH of 4.8 and 4.8 respectively and TOC concentration of 2.2 and 2.1% respectively. The Transition zone soil is sandy clay loam to clay with depth, however the SS soil is a sandy loam grading to sand with depth.

The X-ray diffraction results reveal that quartz is present in all the soils tested, including the Gleysols SF and SS, which is also evident from sand detected in the hand texture assessments.

TABLE 1 | Full soil profile TOC%, bulk density and soil SOC stock for each sub-environment, parentheses indicate standard error of triplicates.

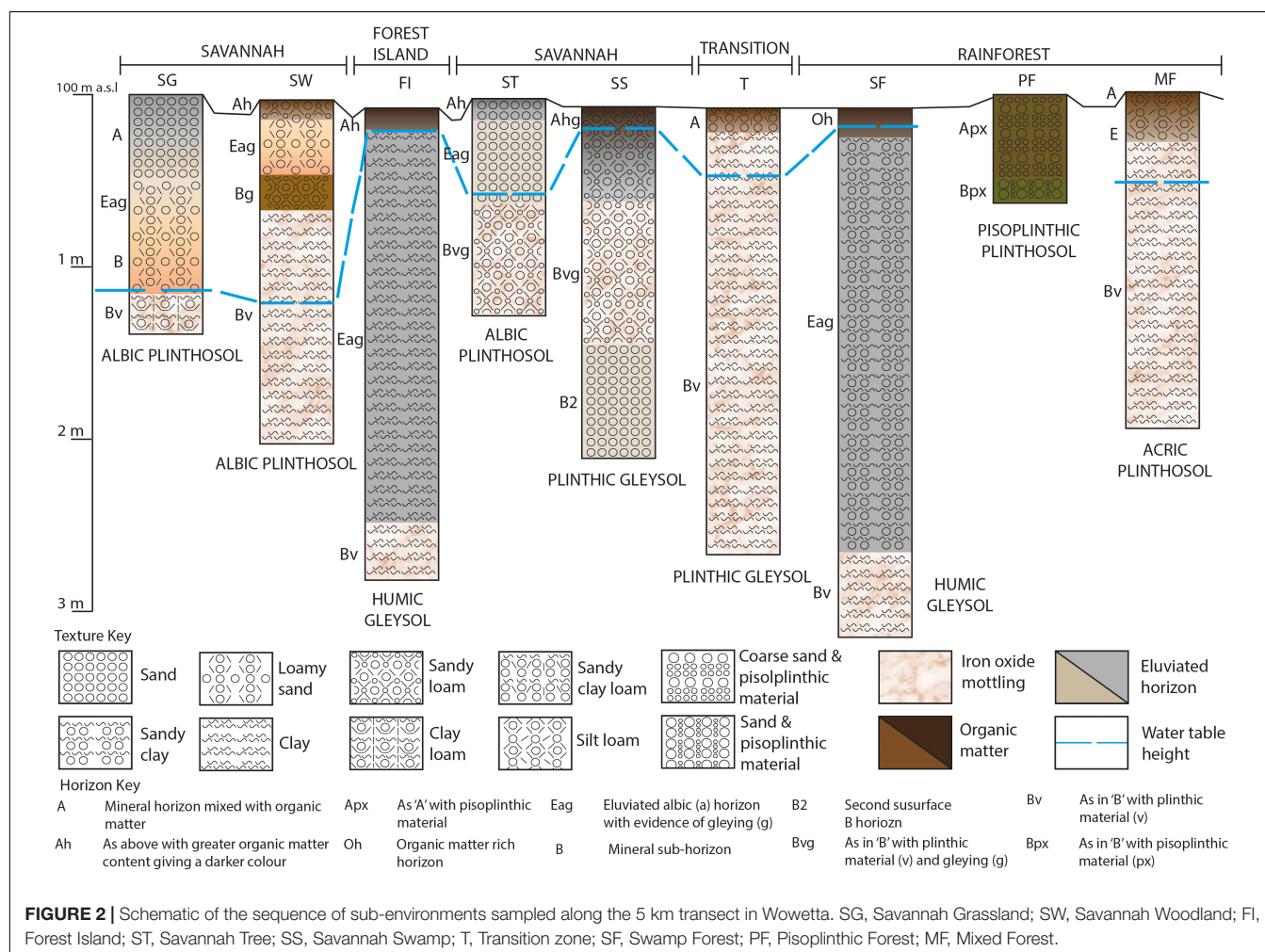
	Boundaries (cm)	TOC (wt%)	D _b (g cm ³)	SOC stock (t C ha ⁻¹)	
Savannah grassland (SG)					
Leaf litter/shoots		44.2			
Wood litter					
0–6 cm	6	0.9	1.7	9.1	(0.2)
A	52	0.5	1.7	43.1	(5)
Eag	103	0.03	2	3.0	(0.9)
B	120	0.02	1.8*	0.6	(1.8)
Bv	137	0.03	1.8*	0.9	(1.8)
Total				56	
Savannah woodland (SW)					
Leaf litter/shoots		45.8			
Wood litter		47.8			
0–6 cm	6	1.6	0.9*	9	(0.5)
Ah	9	1.6	0.9*	13.5	(0.9)
Eag	39	0.2	1.3	17.2	(0.8)
Bg	64	0.5	1.9	18.4	(1.6)
Bv	200	0.04	1.6*	14.4	(0.9)
Total				72	
Forest island (FI)					
Leaf litter/shoots		39.6			
Wood litter		44.6			
0–6 cm	6	7.3	0.7	30.8	(1.3)
Ah	27	5.7	1.3*	200.4	(1.3)
Eag	220	0.1	1.3*	24.3	(1.3)
Bv	280	0.03	1.3*	2.5	(1.3)
Total				258	
Savannah tree (ST)					
Leaf litter/shoots		47.2			
Wood litter		47.3			
0–6 cm	6	1.8	0.9	9.9	(0.03)
Ah	10	1.8	0.9	16.4	(1.8)
Eag	98	1	1.1*	98.3	(0.6)
Bvg	125	0.06	1.5	2.6	(0.06)
Total				127	
Savannah swamp (SS)					
Leaf litter/shoots		41.3			
Wood litter		44.1			
0–6 cm	6	2.6	1.1	17.8	(0.9)
Ahg	54	1.8	1.3	131.7	(2.8)
Bvg	105	0.1	1.2*	3	(1.8)
B2	200	0.1	1.2*	7	(1.8)
Total				159	
Transition zone (T)					
Leaf litter		38.3			
Wood litter		45.4			
0–6 cm	6	2.6	1.1	17.3	(0.3)
A	26	1.6	1.5	64.7	(6.8)
Bv	264	0.1	1.6	21.4	(3.2)
Total				103	
Swamp forest (SF)					
Leaf litter		47.2			
Wood litter		46.3			

(Continued)

TABLE 1 | Continued

	Boundaries (cm)	TOC (wt%)	D _b (g cm ³)	SOC stock (t C ha ⁻¹)	
0–6 cm	6	16.2	0.6	59	(5.6)
Oh	27	8.8	1	228.6	(55.6)
Eag	265	0.3	1.3*	92.1	(20.8)
Bv	333	0.1	1.3*	9.1	(20.8)
Total				389	
Mixed forest (MF)					
Leaf litter		39.5			
Wood litter		45.6			
0–6 cm	6	1.2	1.2	8.5	(0.04)
A	23	1.1	1.5	38.4	(1.6)
E	42	0.1	1.7	3.5	(0.1)
Bv	195	0.1	1.7	20.6	(0.4)
Total				71	

Where applicable, surrogate values for soil horizons are indicated by a *.



The clay mineral kaolinite is present in all samples tested except for the savannah sites ST and SW. The ferric iron phosphate mineral strengite is also present in all the soils tested, except for SF. Anatase, nepheline, and scapolite are also present in

small amounts (see **Supplementary Material** for examples of a savannah and rainforest soils). This mineralogy is unlikely to have a significant effect on the physical protection or complexing of organic carbon, as the quartz, clay mineral kaolinite, ferric

iron strengite and other minerals present have a low reactivity (Lindsay, 1979; Haynes, 1982).

SOC and Volumetric Water Content

The surface 0–6 cm and full profile SOC for each horizon are given in **Table 1**. The highest surface and full soil profile SOC stocks were measured in humic Gleysols; the Swamp Forest (59 t C ha⁻¹ and 389 t C ha⁻¹, respectively), followed by the Forest Island (31 t C ha⁻¹ and 258 t C ha⁻¹, respectively); these sites also have the highest volumetric water contents of 75 and 67%, respectively (**Figure 3**). The other sites, consisting of plinthic Gleysols and albic, acric and pisoplinthic Plinthosols, have surface soil averages of approximately 20 t C ha⁻¹ and 41% volumetric water contents. We observe a significant correlation ($R^2 = 0.9$) between surface SOC and surface volumetric water content (**Figure 3**). The correlation is likely an indication of the texture differences in the soils; with the Gleysols generally consisting of more clay than the Plinthosols, and therefore the ability to hold more water. Similar relationships between soil water content and texture have been evidenced in previous studies (Abrol et al., 1968). Further, the wetter soils occurred at topographical depressions within the landscape where it is likely to accumulate as compared to a slope or hill, therefore topography is also likely to play a role in the greater water content of the Gleysols.

Most soils show a decrease in SOC with depth (SG, SS, FI, T, and SF), however some soils show increases in the deepest horizons (SW, ST, and MF). This is most likely due

to greater horizon thickness in the deepest horizons, as well as slight increases in bulk density observed in most horizons with depth. In most cases, however, TOC concentration decreases with profile depth.

Lignin Phenols

Thermally assisted hydrolysis and methylation in the presence of TMAH of solvent-extracted powdered soil causes SOM degradation to give a series of oxygenated aromatic products dominated by the methylated guaiacyl (G) and syringyl (S) lignin phenols from soils across the study transect. **Figure 4** and **Table 2** give the TMAH thermochemolysis products from all of the soils.

The distribution of the thermochemolysis products from a savannah woodland soil demonstrate that the most prominent products included G1 (1,2-dimethoxybenzene), G2 (3,4-dimethoxytoluene), G4 (3,4-dimethoxybenzaldehyde), G6 (3,4-dimethoxybenzoic acid methyl ester), S1 (trimethoxybenzene), S6 (3,4,5-benzoic acid methyl ester), P18 (*trans* 3-(4-methoxyphenyl)-3-propenoic acid methyl ester), G18 (*trans* 3-(3,4-dimethoxyphenyl)-3-propanoic acid methyl ester) and the C_{16:0} fatty acid methyl ester (C_{16:0} FAME) (**Figure 4**).

(¹³C-TMAH Method) Lambda (Λ)

Lambda (Λ) shows the sum of the amounts of the most dominant lignin phenols (see Methods, THM with TMAH analysis). Summed vegetation and surface soil Λ show little variation between sub-environments (**Figure 5**). The most

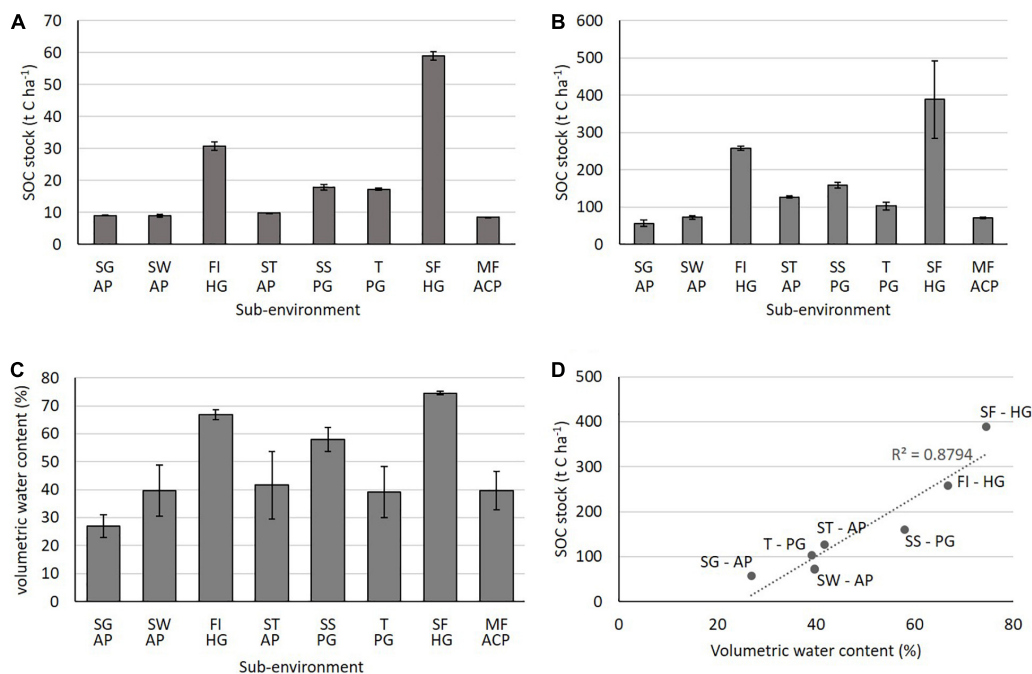


FIGURE 3 | SOC stock at (A) 0–6 cm, (B) total SOC stock of each soil profile (C) average vol. water content for the whole profile of each soil (D) correlation between SOC stock and whole soil profile volumetric water content. SG AP, Savannah Grassland Albic Plinthosol; SW AP, Savannah Woodland Albic Plinthosol; FI HG, Forest Island Humic Gleysol; ST AP, Savannah Tree Albic Plinthosol; SS PG, Savannah Swamp Plinthic Gleysol; T PG, Transition zone Plinthic Gleysol; SF HG, Swamp Forest Humic Gleysol; PF PP, Pisoplinthic Forest Pisoplinthic Plinthosol; MF ACP, Mixed Forest Acric Plinthosol.

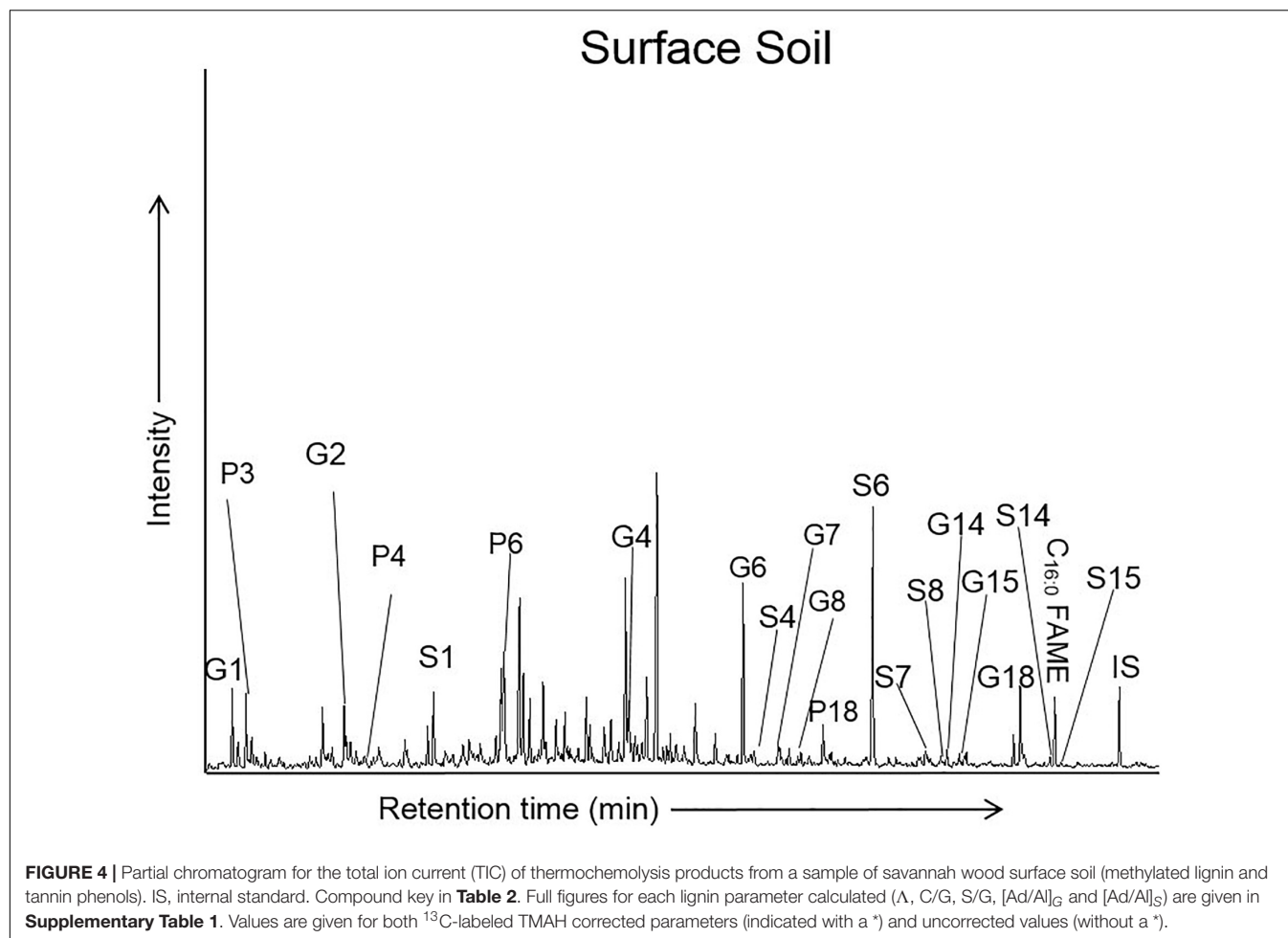


TABLE 2 | Compound codes, names, retention times (RT) and characteristic fragment ions (m/z) of the thermochemolysis products detected across the surface soil samples on the study transect.

Compound code	Compound	RT (min)	m/z
G1	1,2-dimethoxybenzene	35.577	95(50) 123(38) 138(100)
G2	3,4-dimethoxytoluene	43.934	109(45) 137(60) 152(100)
P4	4-methoxybenzaldehyde	45.293	77(40) 135(100) 136(70)
S1	1,2,3-trimethoxybenzene	50.190	110(30) 153(65) 168(100)
P6	4-methoxybenzoic acid methyl ester	55.836	77(15) 135(100) 166(40)
G4	3,4-dimethoxybenzaldehyde	64.64	151(20) 165(50) 166(100)
G6	3,4-dimethoxybenzoic acid methyl ester	73.442	165(90) 181(15) 196(100)
S4	3,4,5-trimethoxybenzaldehyde	74.289	125(30) 181(50) 196(100)
G7	cis 1-(3,4-dimethoxyphenyl)-2-methoxyethylene	76.023	151(35) 179(65) 194(100)
G8	trans 1-(3,4-dimethoxyphenyl)-2-methoxyethylene	76.857	151(35) 179(65) 194(100)
P18	trans 3-(4-methoxyphenyl)-3 propenoic acid methyl ester	79.641	133(35) 161(100) 192(75)
S6	3,4,5-trimethoxybenzoic acid methyl ester	83.046	195(30) 211(50) 226(100)
S7	cis 1-(3,4,5-trimethoxyphenyl)-2-methoxyethylene	86.961	181(20) 209(100) 224(92)
S8	trans 1-(3,4,5-trimethoxyphenyl)-2-methoxyethylene	88.112	181(20) 209(100) 224(92)
G14	threo/ethryo 1-(3,4-dimethoxyphenyl)-1,2,3-trimethoxypropane	88.641	166(15) 181(100) 270(5)
G15	threo/ethryo 1-(3,4-dimethoxyphenyl)-1,2,3-trimethoxypropane	89.608	166(15) 181(100) 270(5)
G18	trans 3-(3,4-dimethoxyphenyl)-3-propanoic acid methyl ester	94.008	191(45) 207(15) 222(100)
S14	threo/ethryo 1-(3,4,5-trimethoxyphenyl)-1,2,3-trimethoxybenzene	96.247	181(10) 221(100) 300(8)
S15	threo/ethryo 1-(3,4,5-trimethoxyphenyl)-1,2,3-trimethoxybenzene	97.094	181(10) 221(100) 300(8)

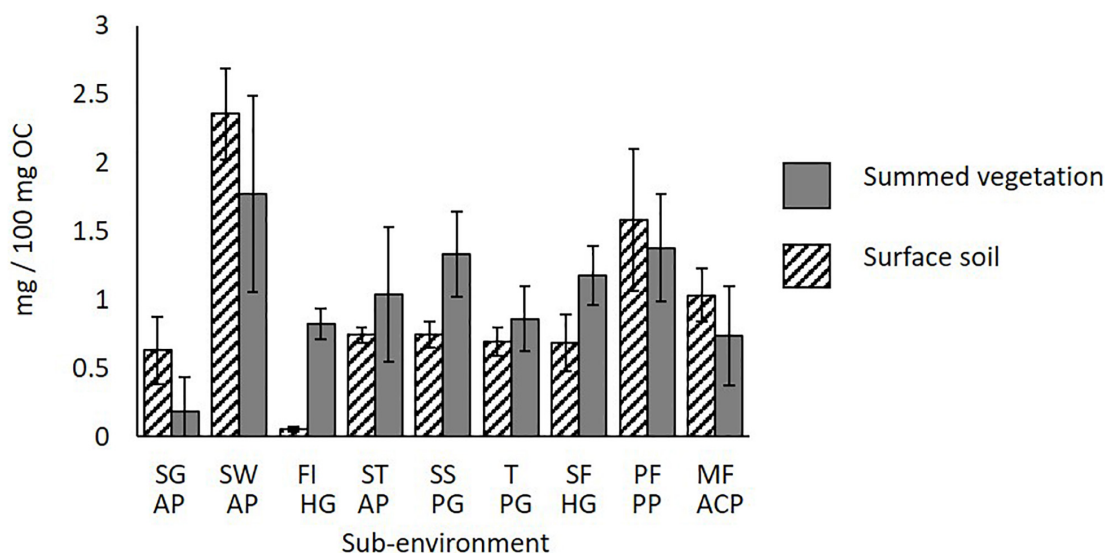


FIGURE 5 | ^{13}C -TMAH corrected Δ values (mg/100 mg OC) for surface soils, and summed inputs = fresh shoot/leaf litter and wood litter across the transect. S.E. bars of 3 replicates shown. SG AP, Savannah Grassland Albic Plinthosol; SW AP, Savannah Woodland Albic Plinthosol; FI HG, Forest Island Humic Gleysol; ST AP, Savannah Tree Albic Plinthosol; SS PG, Savannah Swamp Plinthic Gleysol; T PG, Transition zone Plinthic Gleysol; SF HG, Swamp Forest Humic Gleysol; PF PP, Pisoplinthic Forest Pisoplinthic Plinthosol; MF ACP, Mixed Forest Acric Plinthosol.

notable difference is in the surface soil of the Forest Island, with very low Δ (0.1 mg/100 mg OC) as only lignin phenols G6, S6 and G18 were detected in the THM products. The remainder of the main lignin phenols (G4, G5, G6, S4, S5, P18) have therefore been degraded in the soil, demonstrating their lability.

Further, lignin phenols (specifically G4 and S4) are also shown to be labile in soils of the Savannah Tree, Savannah Swamp, Swamp Forest and Mixed Forest, as the amounts of these phenols in the soils are lower compared to their respective vegetation inputs (see **Supplementary Table 2**). However, in the case of the Savannah Tree and Mixed Forest, the significant difference between vegetation and soils does not affect the overall Δ , as this remains similar.

Greatest Δ was seen in the surface soil of the Savannah Woodland site (2.4 mg/100 mg OC), due to high amounts of phenols G6 and G18 (1.1 and 0.6 mg/100 mg OC, respectively). In comparison, all other Δ values of surface soils across the transect were ≤ 1.6 mg/100 mg OC, with the majority between 0.6 and 0.7 mg/100 mg OC.

In the remaining sites (Savannah Tree, Transition zone, Mixed Forest and Pisoplinthic Forest) soil Δ is similar to the respective vegetation Δ , as phenols G6 and G18 are not significantly degraded. This shows that phenols in these soils are less labile, or phenolic replacement from vegetation is high. Further, the Savannah Grassland surface soil has a slightly greater Δ than vegetation inputs (a difference of 0.4 mg/100 mg OC), suggesting preservation of phenols G18 and G6.

Source Indicators – C/G and S/G

Across the savannah-rainforest transect, the sub-environment surface soils differ in woody/non-woody (C/G) composition; a value above 1 is a non-woody signal and below that is a

woody signal (**Figure 6A**). As anticipated, non-woody ratios are predominant in the Savannah Grassland. Non-woody ratios are also predominant in the Savannah Swamp, Transition zone and Mixed Forest areas. In the latter four sites, selective degradation of guaiacyl over cinnamyl is apparent. However, in Savannah Woodland, Savannah Tree, Forest Island, Swamp Forest and Pisoplinthic Forest soils the opposite trend is apparent, emphasizing selective cinnamyl degradation in surface soils. All wood litter and fresh shoot/leaf litter samples have ratios signaling woody and non-woody input respectively, as expected (Nierop and Filley, 2007).

For the S/G ratio, a value above 1 is predominantly a gymnosperm signal, and below that an angiosperm signal. Except for the Mixed Forest soil, all surface soil S/G values are lower than those for vegetation (<1) showing the loss of the angiosperm signals via degradation (**Figure 6B**). As described in the following section, the Forest Island and Swamp Forest soils show the greatest syringyl degradation.

Level of Oxidation – $[\text{Ad}/\text{Al}]_G$ and $[\text{Ad}/\text{Al}]_S$

Increasing $[\text{Ad}/\text{Al}]_{G/S}$ values indicate increasing oxidation, therefore, it is usually expected that these ratios are higher in soils than vegetation (**Supplementary Table 1** and **Figures 6C,D**; Filley et al., 1999). Oxidation signifies the degradation of lignin phenols. Oxidation ratios $[\text{Ad}/\text{Al}]_{G/S}$ could not be calculated for the surface soil of the Forest Island, as both G4 and S4 were not detected, suggesting an advanced rate of degradation at this location. The Swamp Forest also has a very degraded soil signature – it has the highest syringyl oxidation level in both surface soil and leaf litter (14.1 and 4.79, respectively). Due to the greater bulk SOC stocks in the Gleysols of the Forest Island and Swamp Forest, it was not expected that lignin would be

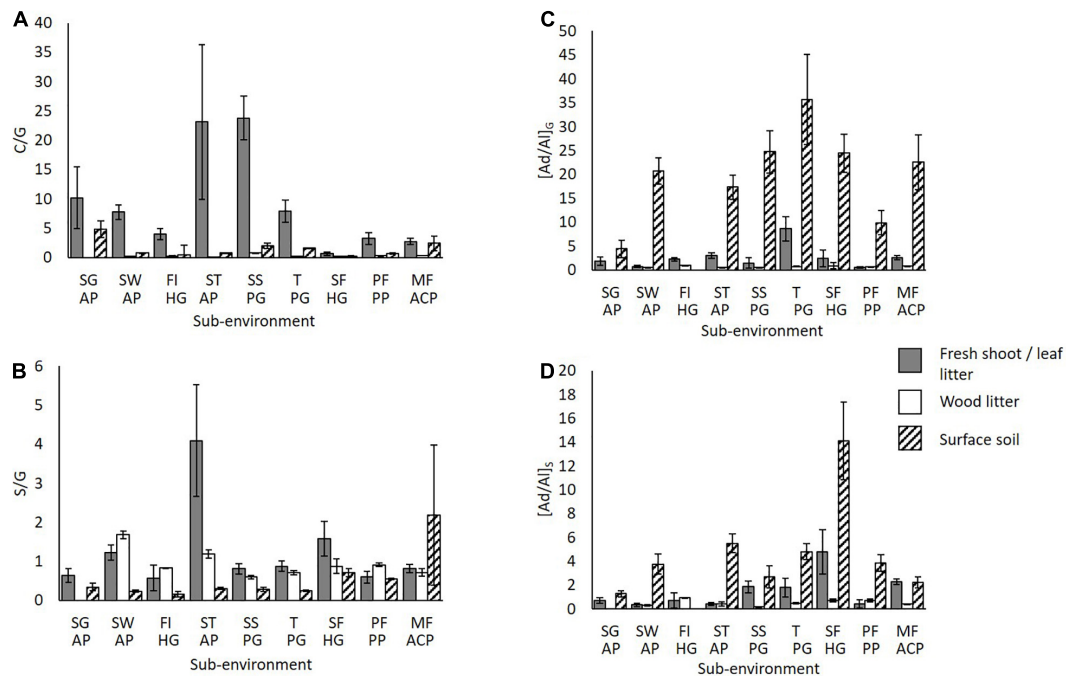


FIGURE 6 | Lignin parameters (A) C/G and (B) S/G of surface soils and vegetation inputs across the transect and oxidation extents of (C) [Ad/Al]_G and (D) [Ad/Al]_S of surface soils and vegetation inputs in sub-environments across the transect. S.E. bars of 3 replicates are shown. SG AP, Savannah Grassland Albic Plinthosol; SW AP, Savannah Woodland Albic Plinthosol; FI HG, Forest Island Humic Gleysol; ST AP, Savannah Tree Albic Plinthosol; SS PG, Savannah Swamp Plinthic Gleysol; T PG, Transition zone Plinthic Gleysol; SF HG, Swamp Forest Humic Gleysol; PF PP, Pisoplinthic Forest Pisoplinthic Plinthosol; MF ACP, Mixed Forest Acric Plinthosol.

degraded to this extent. The Savannah Grassland and Pisoplinthic Forest have the lowest guaiacyl oxidation and the Mixed Forest the lowest syringyl oxidation. All other sub-environment surface soils have comparable oxidation levels.

Wood litter in the study area are less oxidized than soils, as expected, and generally have very low values indicating hardwood [Ad/Al]_G: 0.52–0.92 and [Ad/Al]_S: 0.17–0.75 (Nierop and Filley, 2007). Some surface soils show a similar degree of oxidation to the respective fresh shoot or leaf litter samples, possibly due to rapid polyphenol input from fresh grass shoots, for example the guaiacyl compounds in the soil of the Savannah Grassland (Nierop and Filley, 2007). Similar oxidation degrees between soils and vegetation may also occur due to a high degree of lignin degradation in decomposing leaf litter, such as in the Mixed Forest and Transition zone sites, e.g., [Ad/Al]_G 2.31 and 2.24, and [Ad/Al]_S 2.44 and 2.31 for soil and leaf litter respectively.

Gallic Acid (Tannin) and Demethylated Lignin Sources

Figure 7 shows the amounts of intact lignin, demethylated lignin and gallic acid in syringyl phenol S6 in wood litter (a), fresh shoots/leaf litter (b) and surface soils (c) across the sampling transect. This syringyl phenol showed the strongest alteration after correction with ¹³C-labeled TMAH.

It is evident that gallic acid is most abundant in the Savannah Woodland and Tree areas; vegetation input

averages are 2.6 mg/100 mg OC and 2.16 mg/100 mg OC respectively, and surface soil averages are 1.48 mg/100 mg OC and 0.47 mg/100 mg OC, respectively (Figures 7A–C). The other sub-environments mostly show much lower amounts (<0.27 mg/100 mg OC). Hättenschwiler and Jørgensen (2010) demonstrated that different plant species can have high variation in foliage and leaf litter chemistry within the same rainforest ecosystem, thus, the differences observed here between sub-environments may be expected.

DISCUSSION

Natural Heterogeneity of SOC in Savannah-Rainforest Boundaries

Our quantitative estimates of SOC stocks (Table 1 and Figures 3A,B) along the transect studied align with measurements from other regions of savannah-rainforest mosaics (Batjes and Dijkshoorn, 1999; Schwartz and Namri, 2002), confirming the small-scale natural heterogeneity of soils along vegetation gradients. For example, Plinthosol average surface soil (0–6 cm) SOC stock is significantly lower than that of the Gleysol average; 9 t C ha⁻¹ compared to 31 t C ha⁻¹. Full soil profile averages of Plinthosols are also lower than those of Gleysols (82 t C ha⁻¹ compared to 227 t C ha⁻¹). This difference in SOC stocks is consistent with surface and full profile Gleysol and Plinthosol SOC stock estimates reported for the Brazilian

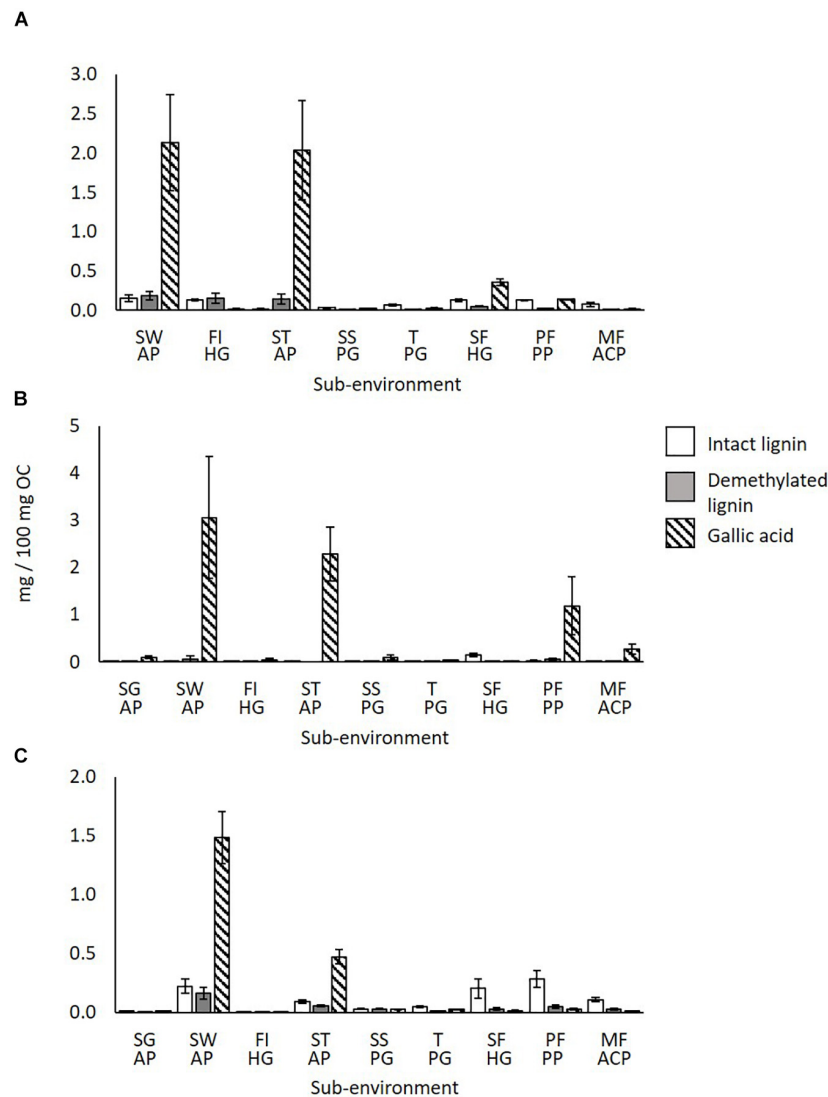


FIGURE 7 | Intact lignin, demethylated lignin and gallic acid amounts across the sampling transect (SG-MF) in **(A)** wood litter, **(B)** fresh shoot/leaf litter, and **(C)** surface soils. S.E. bars of 3 replicates shown. SG AP, Savannah Grassland Albic Plinthosol; SW AP, Savannah Woodland Albic Plinthosol; FI HG, Forest Island Humic Gleysol; ST AP, Savannah Tree Albic Plinthosol; SS PG, Savannah Swamp Plinthic Gleysol; T PG, Transition zone Plinthic Gleysol; SF HG, Swamp Forest Humic Gleysol; PF PP, Pisoplinthic Forest Pisoplinthic Plinthosol; MF ACP, Mixed Forest Acric Plinthosol.

Amazon (Batjes and Dijkshoorn, 1999), although the scale of carbon burial differs significantly between regions.

For example, when comparing our Gleysol estimates of 128 t C ha^{-1} at 0–30 cm depth to Batjes and Dijkshoorn's (1999) study on Amazon Gleysols, they are almost double their average of 70 t C ha^{-1} (0–30 cm). However, comparing further afield to the Congolese basin, Schwartz and Namri (2002) estimate greater surface (0–10 cm) SOC stocks of $74\text{--}109 \text{ t C ha}^{-1}$ in 'swamp forest' hydromorphic soils and peat than we present here for Swamp Forest and Forest Island humic Gleysols at an average of 59 t C ha^{-1} (0–10 cm). Despite these differences, Plinthosols in this study average 38 t C ha^{-1} at 0–30 cm depth, consistent with previous estimates of 40 t C ha^{-1} (Batjes and Dijkshoorn, 1999). Schwartz and Namri (2002) show that similar Ferrallitic

savannah soils could have greater stocks; ranging between $15\text{--}30 \text{ t C ha}^{-1}$ at 0–10 cm depth in their Congolese study region. Together these studies from different locations emphasize the broad range and diversity of soil carbon stocks at the small scale. Each study therefore represents one facet of a larger and not yet well constrained carbon puzzle that characterizes forest-savannah boundaries.

Soil organic carbon variation across different soil and vegetation types has been explained in various studies by a textural and hydromorphic effect, where high water content aligns with high SOC stocks through the anoxic conditions and subsequent preservation of carbon (Schwartz and Namri, 2002; Abbott et al., 2013). SOC variation across the transect in Guyana supports this general relationship as humic Gleysols have

both the highest SOC stock and volumetric water content. The plinthic Gleysol under Savannah Swamp shows a relatively strong hydromorphic effect on both surface and full soil profile SOC, which are both higher than the Plinthosols. The hydromorphic effect on the plinthic Gleysol in the Transition zone is harder to document, with the surface SOC stock being greater than in the Plinthosol, yet the volumetric water content being similar. The same challenge applies to albic Plinthosols; Savannah Woodland and Savannah Tree soils have higher volumetric water contents than Savannah Grassland, but their surface SOC stocks are similar. As stated in the methods, water content was only measured at one point in time, and therefore an annual average may make this relationship clearer, but was beyond the scope of the study. However, the relationship indicates that texture is likely to be an important factor, as demonstrated by the Gleysols which have a predominantly clay texture likely contributing to greater water content, and Plinthosols which have a predominant texture of sand, and are thus drier. Previous research has also evidenced a relationship between texture and water content (Abrol et al., 1968). Therefore, predominantly clayey soils with greater water contents (swamp savannah and forest sites) show the hydromorphic effect more clearly than those with lower water contents (transition zone and savannah sites). Topography also likely influences water content, as Gleysols were found in depressions within the landscape, as previously noted.

The direct vegetation effect on SOC can be assessed between Savannah Grassland and Savannah with trees, as both contain sandy soils, which are otherwise absent in the forest environments studied. Comparing the two, it may be that the Savannah Woodland and Tree sites have higher vegetation inputs and therefore greater SOC stocks than the Savannah Grassland.

However, grasses can have a significant input of SOC through their roots, therefore it is possible that SOC stocks between these different savannah sub-environments may be similar (Lal, 2004; Stahl et al., 2016). The SOC results from the study transect do show that carbon stocks are similar at the surface. We therefore conclude that the vegetation effect in savannah environments is small, at least on the transect analyzed, supporting previous studies in the Congo by Schwartz and Namri (2002). Abiotic factors such as the predominant texture of sand is thus likely to be more influential than vegetation; contributing to the low SOC stocks of these savannah environments through both the poor ability of carbon to be protected and the effect of leaching of carbon easily through sand particles after the wet season flooding.

Lignin Degradation Controls and Their Relation to SOC From Rainforest to Savannah

The textural, hydromorphic and topographical effects on soil carbon carries down from the bulk SOC to the molecular carbon scale. This is evidenced by the complete loss of guaiacyl G4 and syringyl S4 in the Forest Island Gleysol, as well as the highest $[Ad/Al]_S$ values in both surface soils and leaf litter of the Swamp Forest site (see Figure 8). Further, the Forest Island has the lowest total lignin phenols (Λ) on the study transect (see Figure 5). The Swamp Forest also has a relatively low Λ compared to other soils on the transect (high lignin oxidation in $[Ad/Al]_{G\&S}$ values also result in decreases in Λ values). This high molecular degradation was not initially anticipated due to the high water content and organic matter-rich nature of the soils observed when sampling.

Figure 8 below demonstrates the difference between SOC stocks and phenolic degradation ($[Ad/Al]_{G\&S}$) and lignin phenol

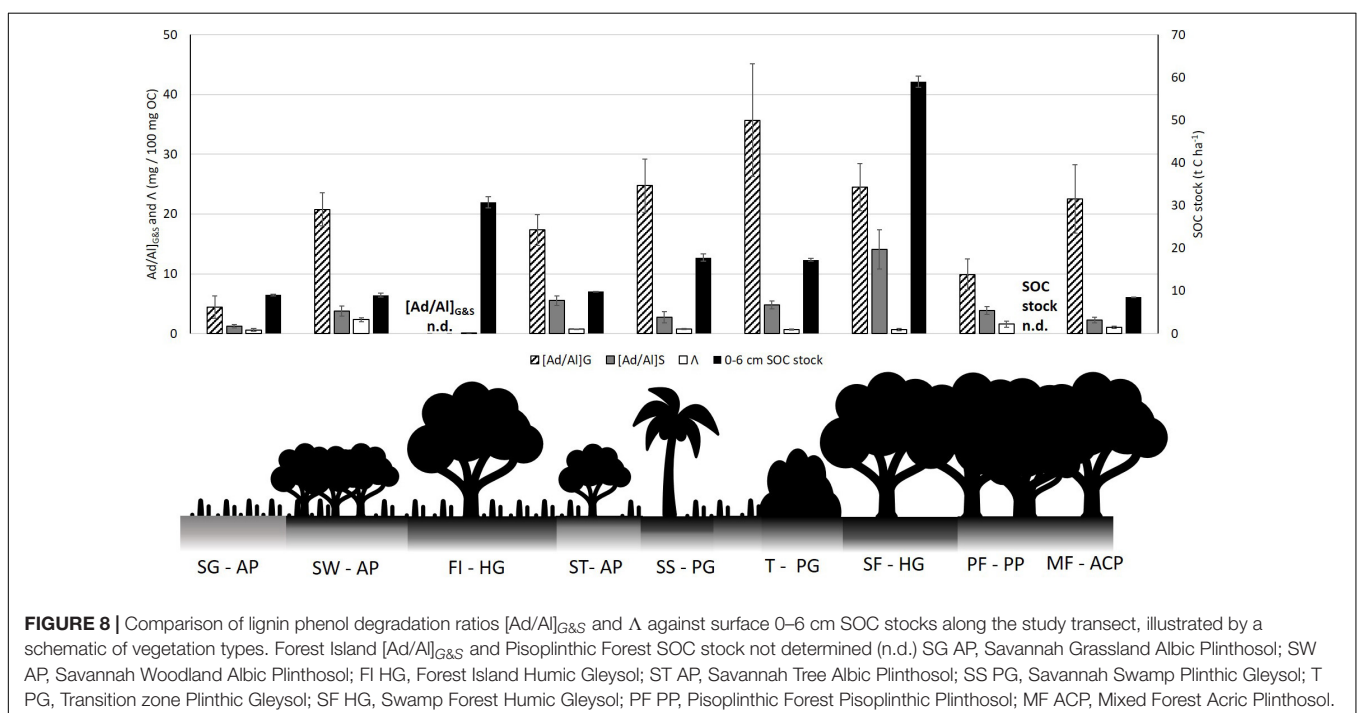


FIGURE 8 | Comparison of lignin phenol degradation ratios $[Ad/Al]_{G\&S}$ and Λ against surface 0–6 cm SOC stocks along the study transect, illustrated by a schematic of vegetation types. Forest Island $[Ad/Al]_{G\&S}$ and Pisoplinthic Forest SOC stock not determined (n.d.) SG AP, Savannah Grassland Albic Plinthosol; SW AP, Savannah Woodland Albic Plinthosol; FI HG, Forest Island Humic Gleysol; ST AP, Savannah Tree Albic Plinthosol; SS PG, Savannah Swamp Plinthic Gleysol; T PG, Transition zone Plinthic Gleysol; SF HG, Swamp Forest Humic Gleysol; PF PP, Pisoplinthic Forest Pisoplinthic Plinthosol; MF ACP, Mixed Forest Acric Plinthosol.

sum (Λ). Where SOC stocks are highest, the degradation extent of lignin guaiacyl and syringyl is high. This is particularly evident with $[Ad/Al]_S$ in the Swamp Forest Gleysol and the lack of phenols in the Forest Island Gleysol. Similarly, Savannah Woodland and Savannah Tree plinthosols show decoupling through the reverse trend – low SOC stocks and high amounts of lignin phenols, as well as high tannin phenol amounts (see **Figure 7**). SOC stocks from Savannah soils through to the Transition soil increase slightly, which is tracked by phenolic degradation. These then decouple again at the Swamp Forest. The decoupling between the degradation of lignin and the SOC stocks contradicts the initial hypothesis stated, that SOM degradation is inversely associated with SOC stock.

Such high degradation of lignin phenols in the Forest Island and Swamp Forest is likely due to the physio-chemical environment and biota surrounding the lignin, which allows for degradation processes to occur, for example the clay texture leading to a greater soil moisture, allowing them to be mineralized by microbes (see **Figure 6**). The extent of its breakdown suggests that it may be older carbon, although radiocarbon dates would be needed to verify this. The greater clay content of these rainforest soils may hold onto moisture for a longer period than the sandier soils, potentially favoring microbial communities to be active in degrading lignin for more extended periods of time. As previously noted, the volumetric water content was measured from the soil samples collected and therefore only represent one point in time. However, this variable is likely to fluctuate temporally on a seasonal and longer time scale, which will impact microbial activity and rates of organic matter decomposition. Indeed, Bovolo et al. (2012) evidence the seasonality of rainfall more widely across Guyana, showing the two wet and dry seasons of the rainforest, and one wet and dry season of the savannah. In this study it was not possible to assess annual variation in soils and how such seasonality therefore affects SOC, however the effect of weather patterns on soils has been noted by other studies. For example, annual variation in precipitation and short-scale rain events after and between dry periods may break open physically protected OM and activate microbial populations, allowing them to release phenol oxidase for decomposition (Austin et al., 2004; Fenner and Freeman, 2011; Abbott et al., 2013; Schellekens et al., 2015a,b,c). Therefore, lignin which has been previously protected in soil aggregates may become exposed to degradation through wet-dry cycles.

Further, fine-grained soils can protect organic carbon to a greater extent than coarse-grains, as organic matter turnover is governed by accessibility not recalcitrance (Dungait et al., 2012). Incorporation into clay minerals and formation of organo-mineral complexes with the iron oxides present in the soils from underlying plinthic material (**Figure 2**) may therefore play a role in protecting the SOC. However, significant incorporation into clay minerals is unlikely given that the X-ray diffraction results show that quartz and kaolinite constitute the mineralogy of all the soils sampled along the transect (see **Supplementary Figures 1, 2** for examples of a savannah and rainforest soil mineralogy). The presence of the ferric iron phosphate mineral strengite detected in the soils also suggests a low ability for

organic matter to complex, as this mineral is known to be stable at low pH in soil (Lindsay, 1979; Haynes, 1982). Conversely, iron oxy-hydroxides (such as hematite) were not detected, which may have otherwise induced more significant organo-mineral complexing (Gu et al., 1994; Sollins et al., 1996; Arnarson and Keil, 2000; Lützow et al., 2006). Therefore, it is likely that the moisture retentive properties of the clay textured soils and their topographical position play a more important role in the degradation of lignin.

The Forest Island lies surrounded by savannah and has much more exposed soil than, for example, the Swamp Forest. Patches of Forest Island on the boundary may therefore undergo more pronounced dehydration during the dry season as well as drying out faster following wet periods, allowing decomposition for a longer portion of the year. This exposure may explain the more significant phenolic depletion in comparison to the Swamp Forest soil, resulting in greater carbon degradation during both the dry and the wet season.

A previous study evidences leaching of dissolved organic matter and humic substances, including lignin, from forest soils into river waters during rainfall events close to the study site at Iwokrama in Central Guyana (Pereira et al., 2014). Leaching is therefore more than likely to account for some forest soil lignin decomposition. This may be the case particularly at Mixed and Pisoplinthic Forest sites, where lignin oxidation is apparent and the water content and SOC stock of the former site is low.

As the transect grades from forest to savannah, the high level of guaiacyl degradation (see **Figure 6**) in plinthic Gleysols in the Savannah Swamp can also be explained by a texture and hydromorphic effect. The clay texture of this soil means that it retains a higher level of moisture than Plinthosols in other areas of savannah and rainforest. It is likely that this degraded lignin signature is part of old carbon which has built up to an extent, as it also has a higher SOC stock than other savannah soils.

Unique on the study transect, the Savannah Woodland and Savannah Tree soils have significantly high gallic acid contents, which are reflected in the molecular content of the wood and leaf litter inputs. This is likely further influenced by the relatively low soil moisture which may constrain microbial mineralization. The high gallic acid contents, along with high Λ contents, do not have a significant effect on these SOC stocks however, which are some of the lowest on the transect.

The Savannah Grassland acric Plinthosol has a similar Ad/Al oxidation level between the soil and fresh grass shoots, which indicates fresh lignin inputs here (Nierop and Filley, 2007). This results in a similar Λ to the majority of soils on the transect. Grass shoot Λ is significantly lower than other vegetation inputs on the transect however, which constrains the SOC stock. It is likely that grass roots also input significant amounts of carbon into the soil (Stahl et al., 2016), however the low SOC stock suggests that this is lost due to abiotic or biotic factors. Reinforcing this, after the wet season which causes flooding, it is likely that carbon will be leached from the soil profile as the coarse grains of the sandy soil allows for drainage once heavy rains subside. Leaching of molecular compounds and bulk SOC is likely to have a stronger effect on the sandier

soils of the savannah compared to those with greater clay contents. Under the exposed short grassland cover, soil may also undergo degradation of lignin phenols via photooxidation, resulting in lower SOC stocks (Hernes and Benner, 2003; Gallo et al., 2006; Frouz et al., 2011). The soil water content for Savannah Grassland, at the time of sampling in the dry season, was 27%, therefore indicating that some microbial mineralization of carbon may also be possible; although this will likely be more constrained than soils with a greater water content (Austin et al., 2004).

Implications for Future Savannah-Rainforest Boundary Scenarios

This study has revealed that the texture of soils and the water content related to this creates different physio-chemical environments and systems of carbon degradation. In Gleysols, high SOC contents and degraded lignin phenols indicate that carbon may persist and build up over time, being slowly degraded, whereas in Plinthosols, low SOC contents may indicate that carbon does not persist to the same extent. Here, the less degraded lignin phenols indicate fresh lignin inputs (Nierop and Filley, 2007).

As has been discussed above, the various sub-environments along this continuum bear features and mixtures of both carbon cycling systems, which will respond to land use or climate change through altered carbon inputs and soil moisture levels. Climate predictions and El Niño Southern Oscillation (ENSO) effects suggest a shift to higher temperatures (an annual average increase between 0.5–3°C) across northern South America (IPCC, 2013, 2014). More localized models of the Guiana Shield region (partly covering northern Brazil, southern Venezuela, eastern Colombia and the whole of Guyana, Suriname and French Guiana) also indicate increased temperatures (~2°C) and variable changes in precipitation. These changes likely give rise to longer dry seasons, heavier rainfall events in wet seasons but also increased run off and evapotranspiration rates (Bovolo et al., 2018). Whilst the latter study provides predictions on a more regional scale than the IPCC, there may be further localized differences on and surrounding the transect itself, therefore there are limitations to applying these predictions to our data. In such a scenario, rainforest-dieback is likely to occur, causing a gradual transition to more widespread savannah landscapes and a low carbon input system. Lal (2005), Cramer et al. (2004), Don et al. (2011), and Bonini et al. (2018) have shown that when climate change or forestry activity causes this principal shift in vegetation, SOC reduces. Our study provides evidence that relics of previous Swamp Forests are able to remain as Forest Islands with high SOC stocks, likely due to their resistance, stemming from their position in topographical depressions, clayey and therefore wetter soils. However, the presence of highly degraded lignin phenols in these Forest Island soils reveal their fragility to change if perturbed any further. We therefore consider that the carbon pool of these Forest Islands surrounded by savannah is likely to become more decomposed under warmer and drier conditions. The Savannah Swamp and

Transition zone Gleysols illustrate that even in topographical depressions with wetter soils than other savannah sites, SOC stock can be significantly lower; therefore, if the Forest Island degrades to similar vegetation, it may lose some of its SOC stock. The magnitude of a decrease in SOC stocks is difficult to gauge based on the data and due to the yet unknown potential change in soil water content and full understanding of soil carbon dynamics. We therefore suggest further investigation into such dynamics to estimate the potential losses of SOC on this transect.

CONCLUSION

We find that parent material, topography, texture and hydrological conditions at each sub-environment are very influential upon SOC stock. Clay-textured soils with the highest moisture content (Gleysols) have the highest SOC stocks, particularly those under forest vegetation in depressions (swamp and island forests). The Savannah swamp Gleysol has a substantial carbon stock but it is significantly lower than those of the Swamp Forest and Forest Island. Plinthosols have significantly lower SOC stocks than Gleysols, regardless of savannah or rainforest vegetation. Therefore, soil type is a more accurate indicator of SOC stock than vegetation type.

Analysis of the SOM characteristics in the surface soils of the sub-environments reveals clear differences in molecular composition. The Savannah Woodland Plinthosol has the greatest amount of lignin and tannin products and thus most unaltered plant derived material (although some degradation is apparent). Savannah Grassland and Mixed Forest Plinthosols have lower amounts of lignin and tannin products, although the type of phenols that are present indicate that they are fresher inputs. Despite Forest Island and Swamp Forest Gleysols having the highest SOC stocks, these soils have some of the most degraded lignin and tannin amounts across the transect. The original hypothesis is therefore contradicted by our results, in that SOM degradation is inversely related to SOC stock. The results suggest that the physio-chemical environment of the soil surrounding the lignin phenols dictates their degradation extent: in Gleysols lignin degrades slowly over time, as bulk carbon stocks accumulate; whereas in Plinthosols, we observe fresher inputs of lignin phenols and lower SOC stocks.

Upon deforestation due to land use or climate change, as predicted for northern South America and the Guianas more regionally, the high lignin oxidation and thus molecular carbon degradation of Gleysols indicates a vulnerability to savannah encroachment. Forest Islands isolated from the main forest biome are the most vulnerable to change, and could lose some of their SOC stocks. In order to better elucidate the potential loss of SOC and further understand the soil molecular dynamics, we suggest areas for further exploration below.

Recommendations for Future Work

This study has set the framework for further investigations into soil carbon dynamics on the rainforest-savannah boundary

in Guyana, by assessing both bulk and molecular soil organic carbon. The results have highlighted important areas for future study in order to better understand the dynamics of SOC, which is of significance regarding climate change. We therefore suggest that this consists of:

1. Temporal studies of soil moisture, bulk and molecular SOC in order to better elucidate the relationship between these variables.
2. Radiocarbon dating of SOC in order to clarify its age and better understand turnover dynamics.
3. Analysis of lignin with depth in the soil profiles to gain further information on its fate in soil.
4. Analysis of the soil microbial communities to explore their effect on SOC dynamics.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

JB planned and carried out the field work, laboratory work, data analysis, and wrote the manuscript. The research plan was conceived by JB, GA, and TW. GA and TW supervised the project and gave suggestions for the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/ffgc.2021.629600/full#supplementary-material>

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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